This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representation of The original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.

THIS PAGE BLANK (USPTO)



WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) (51) International Patent Classification 5: WO 93/12275 (11) International Publication Number: A1 D01F 11/02, 2/28, A61L 15/28 (43) International Publication Date: 24 June 1993 (24.06.93) (74) Agent: HALE, Stephen, Geoffrey; J.Y. & G.W. Johnson, Furnival House, 14-18 High Holborn, London WCIV (21) International Application Number: PCT/GB92/02283 6DE (GB). 9 December 1992 (09.12.92) (22) International Filing Date: (81) Designated States: AT, AU, BB, BG, BR, CA, CH, CS, DE, DK, ES, FI, GB, HU, JP, KP, KR, LK, LU, MG, MN, MW, NL, NO, NZ, PL, RO, RU, SD, SE, US, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, SN, TD, TG). (30) Priority data: 10 December 1991 (10.12.91) GB 9126193.3 (71) Applicant (for all designated States except US): COURTAULDS PLC [GB/GB]; 50 George Street, London W1A 2BB (GB). (72) Inventors; and
(75) Inventors/Applicants (for US only): BAHIA, Hardev, Singh [GB/GB]; 18 St. Lukes Road, Coventry CV6 4JA (GB).
JAMES, Jim, Robert [GB/GB]; 60 Ambleside Road, Bedworth, Warwickshire CV12 8RR (GB). **Published** With international search report.

(54) Title: CELLULOSIC FIBRES

(57) Abstract

Carboxymethyl cellulose fibre having a degree of substitution of at least 0.1, preferably 0.2-0.5, carboxymethyl group per glucose unit, is derived from solvent-spun cellulose fibre, for example by reaction with a strong alkali and a monochloroacetate reagent. The fibre has an absorbency of at least 8 grams, usually at least 15 grams, 0.9 % saline solution per gram of fibre and a tenacity of at least 10, usually at least 15, cN/Tex. It can be used for absorbent personal products.

P

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

				MR	Mauritania
AT	Austria	FR	France	MW	Malawi
AU .	Australia	GA	Gabon	NL	Netherlands
BB	Barbados	GB	United Kingdom		
BE	Belgium	GN	Guinca	NO	Norway
	Burkina Faso	GR.	Greece	NZ	New Zealand
BF		ผบ	Hungary	PL	Poland
BC	Bulgaria	IE.	Ireland	PT	Portugal
BJ	Benin	iT.	Italy	RO	Romania
BR	Brazii .		-	RU	Russian Federation
CA	Canada	JP.	Japan	SD	Sudan
CF	Central African Republic	*KP	Democratic People's Republic	SE	Sweden
CG	(ongo		of Korca	_	Slovak Republic
CH	Switzerland	KR	Republic of Korea	sk	
CI	Cote d'Ivoire	K2	Kazakhstan	SN	Senegal
	Cameroon	1.1	Liechtenstein	รบ	Soviet Union
CM		LK	Sri Lanka	TD	Chad
cs	Czechoslovakia	1.0	Luxembourg	TG	Togo
CZ	Czech Republic			UA	Ukraine
DΕ	Ciermany	MC	Monaco	us	United States of America
DK	Denmark	MC	Madagascar	VN	Viet Nam ,
ES	Spain	Ml.	Mali	V14	4 the 1 there 4
FI	Finland	MN	Mongolia		<u>•</u>

- 1 -

CELLULOSIC FIBRES

Field of the invention

This invention relates to cellulosic fibres having high absorbency for aqueous fluids. Such fibres are used in 5 absorbent personal products, such as disposable diapers, tampons, sanitary napkins and incontinence pads, and in absorbent wipes, and may also be used as part of the fibre content in moisture-absorbing clothing or in towels. The absorbency of cellulosic fibres can be increased by 10 incorporating highly absorbent chemicals in the fibres or by the cellulose itself. modification of incorporation of highly absorbent chemicals has the risk that the chemicals may be released from the fibres. The present invention is concerned with chemically modified 15 cellulosic fibres, in particular carboxymethylated cellulose fibres.

Background art

Carboxymethyl cellulose in powder form is well known 20 commercially as a thickener. It is produced by reaction of cellulose pulp with a strong alkali such as sodium hydroxide and monochloroacetic acid or a salt thereof. There have been several suggestions for the production of carboxymethyl cellulose fibres but these have not been widely used production of describing the Documents 25 commercially. GB-A-2220881, GB-Acarboxymethyl cellulose fibres are 2094802, US-A-3731680, US-A-1736714, JP-A-49-55993, JP-A-56-15458, JP-A-3-825 and JP-A-3-269144, and articles in J. Applied Polymer Science, volume 17 (1973) at pages 3375-3388 30 and Textile Research J., 1971, pages 680-685. The references describe the production of carboxymethyl cellulose fibres from regenerated cellulose (viscose rayon) fibres or from cotton. There are problems in achieving highly absorbent fibres which are free of surface stickiness and which are 35 strong enough to be processed on textile machinery.

- 2 -

Disclosure of invention

We have found according to the invention that carboxymethyl cellulose fibre of greater absorbency and strength free from surface stickiness can be produced from 5 solvent-spun cellulose fibre.

Carboxymethyl cellulose fibre according to the invention has a degree of substitution of at least 0.1 carboxymethyl group per glucose unit and is characterised in that the fibre is derived from solvent-spun cellulose fibre and has an absorbency of at least 8 grams 0.9% saline solution per gram of fibre, as measured by the free swell method, and a tenacity of at least 10 cN/tex.

A process according to the invention for the production of carboxymethyl cellulose fibre by reacting 15 cellulose fibre with a strong alkali and a monochloroacetic reagent selected from monochloroacetic acid and salts thereof is characterised in that the cellulose fibre used is solvent-spun cellulose fibre.

Solvent-spun cellulose fibres are fibres spun from a opposed to solvent, as 20 solution of cellulose in a regenerated cellulose fibres which are spun from a solution of a cellulose derivative (cellulose xanthate) which is reconverted to cellulose in the bath into which the fibres are spun. Examples of solvents for cellulose are tertiary amine formamide/nitrogen tetroxide N, N-dimethyl 25 N-oxides, mixtures, dimethyl sulphoxide/paraformaldehyde mixtures and solutions of lithium chloride in N,N-dimethyl acetamide or N-methyl pyrrolidone. The preferred solvents for use in producing solvent-spun cellulose fibres are tertiary amine 30 N-oxides. The production of solvent-spun cellulose fibres is described for example in US-A-4246221 and US-A-4196281, which give examples of preferred tertiary amine N-oxides. The solution of cellulose is spun through an air gap into a bath of a non-solvent for cellulose, usually water, where

- 3 -

the cellulose is precipitated in fibre form.

Solvent-spun cellulose fibre has two main differences in structure from regenerated cellulose fibre and cotton. It has a substantially uniform structure across its cross-section and has greater crystallinity. Regenerated cellulose and cotton fibres both have a structure which includes a relatively dense skin at the surface of the fibre. Solvent-spun cellulose fibre has no such skin. We believe that either or both of these properties is important in producing carboxymethyl cellulose fibres of high absorbency without weakening the fibre during the carboxymethylation process to such an extent that it loses its fibre structure.

When carrying out the process of the invention the alkali and the monochloracetic reagent can be applied to the sequentially. 15 cellulose fibre simultaneously or cellulose fibre can be in the form of a tow, yarn, staple fibre or fabric, for example a woven, knitted or non-woven fabric. If a non-woven fabric is used, it is preferably one in which the fibres are relatively securely bound in the 20 fabric, for example a hydroentangled or needled fabric. The yarn, tow or fibre can be a blend of the cellulose fibre with one or more other fibres such as polyester or nylon which are unaffected by the carboxymethylation process. the case of tow, yarn or staple fibre the fibre can be dry 25 fibre as commercially sold or it can be never-dried fibre, is fibre which has not been dried after fibre formation. The rate of uptake of reagents by the fibre and the rate of reaction with the cellulose of the fibre may be somewhat faster using never-dried fibre. If never-dried 30 fibre is used, its water content is preferably controlled to be less than 150%, for example 50 to 100%, by weight by mangling if necessary.

The alkali and the monochloroacetic compound are preferably applied from aqueous solution. The alkali is preferably an alkali metal hydroxide such as sodium

- 4 -

hydroxide or potassium hydroxide and is preferably used at a concentration of at least 2% by weight, more preferably 4% or 5% or more, up to 15% by weight, more preferably up to 10%. The monochloroacetic acid is preferably used in salt 5 form, usually the salt corresponding to the alkali used, for example sodium monochloroacetate with sodium hydroxide. The preferably used is monochloroacetic reagent concentration of at least 5% by weight, more preferably at least 10%, up to 35% by weight, more preferably up to 25%. 10 The alkali and monochloroacetate salt are preferably used in approximately equimolar amounts, for example at a molar hydroxide and 0.8-1.2:1. Sodium ratio monochloroacetate are preferably used at a weight ratio of 1:2.5-3.5, most preferably about 1:2.9. If monochloroacetic 15 acid is used, the molar ratio of alkali to monochloroacetic acid is preferably about 2:1.

The alkali and the monochloroacetic reagent can be applied from solution in a mixture of water and a polar organic solvent. For example, sodium hydroxide can be 20 dissolved in water at up to 35% by weight and sodium monochloroacetate dissolved in water at up to 45% by weight, and the solutions can be mixed and diluted with an alcohol such as ethanol or industrial methylated spirits to give the required concentration of reagents in the aqueous organic solvent mixture.

Various procedures can be used for applying the alkali and monochloroacetic reagents to the fibre. The fibre can be immersed in a solution of a reagent at elevated temperature, for example at least 50°C up to the boiling point of the 30 solution. When the fibre is in the form of a continuous textile material such as a tow or a fabric, the reagent can be applied by padding followed by drying at an elevated temperature. The alkali and monochloroacetic reagents can be applied sequentially, in which case it is preferred but not applied before is alkali that the 35 essential monochloroacetic reagent, or they can be applied together in

- 5 -

a single solution. The padding procedure has the advantage that drying can take place at higher temperatures, allowing shorter reaction time. The drying temperature can, for example, be in the range 50 to 200°C. Drying temperatures of 5 above 100°C allow reaction times of for example 2 to 10 minutes compared to 4 to 30 minutes for reactions carried out at below 100°C. The amount of liquid taken up during padding is preferably 50 to 300% by weight; the tow or fabric can be squeezed, for example by mangling, after 10 padding if required to give the desired take-up. Too high a liquid level may mean that the time required for drying is greater than the time required for the carboxymethylation reaction. Although still effective, this is wasteful of energy. Care should be taken to avoid substantial weakening 15 of the cellulose fibre by thermal degradation. The most preferred reaction temperatures may be in the range 80 to 150°C, particularly temperatures of 90°C and above such as 90 to 120°C. The heating of the treated fibres should preferably not be such as to completely dry the fibres, a 20 moisture content of 5 to 20% by weight after drying being preferred. This avoids brittleness of the dried fibres. Heating can take place in an oven or in a hot tunnel dryer in a continuous process.

One example of a procedure for carrying out the process 25 of the invention comprises padding the fibre with alkali, for example aqueous sodium hydroxide, and drying at a temperature above 80°C, followed by treatment with the sodium for example monochloroacetic reagent, monochloroacetate. The alkali-treated fibre can be immersed to 100°C for 15 to 60 30 in an aqueous solution at 50 minutes, followed by drying. Alternatively, the sodium monochloroacetate solution can be applied by padding, followed by a drying step at a temperature above 80°C for 5 to 15 minutes. The fibre in tow or fabric form can be 35 mangled after padding with the sodium hydroxide or after padding with the sodium monochloroacetate, or both, to reduce the liquid take-up on the fibre in each case. If the

- 6 -

fibre is mangled to reduce the liquid take-up, the concentration of the solution used should be increased to give a similar level of reagent on fibre. The order of treatment can be reversed, that is the fibre can be padded with sodium monochloroacetate solution and optionally mangled and dried, followed by padding with sodium hydroxide, optionally mangling and drying. In either case, care should be taken when making up the second treatment solution in a continuous process to allow for reagent from the first treatment solution carried by the fibre into the second treatment solution and re-dissolved.

and hydroxide sodium Alternatively, the applied simultaneously. monochloroacetate can be simultaneous application of sodium hydroxide and sodium 15 monochloroacetate may be preferred because only one drying step is used and the overall reaction time is reduced. solution containing the required concentration of sodium hydroxide and sodium monochloroacetate can be prepared by mixing the solutions of these reagents which have been 20 separately prepared or by dissolving sodium hydroxide in a solution of sodium monochloroacetate. The mixed solution can be applied by immersion or by padding, optionally followed by mangling, and drying at elevated temperature. solution containing both sodium hydroxide and sodium 25 momochloroacetate should preferably not be held for an extended time at an elevated temperature, since reaction of the NaOH and ${\rm ClCH_2COONa}$ to form sodium chloride and sodium glycollate can take place. The sodium hydroxide and sodium monochloroacetate solutions can be mixed just before 30 application to the fibre, or the separate solutions can simultaneously be sprayed onto the fibre, for example by sprays arranged at right angles to each other. If the mixed hydroxide) sodium (eg of alkali solution monochloroacetic reagent (eg sodium monochloroacetate) has 35 to be stored, it is preferably held at a temperature of 20°C or below, for example 0 to 5°C. Storage at 20-40°C of fibre treated with both the alkali and the monochloroacetate is

- 7 -

preferably avoided. It is usually most convenient to heat the fibre immediately after padding to effect the carboxymethylation reaction. Alternatively, the padded fibre can be stored at a temperature below 20°C, preferably in the range 0 to 5°C, before heating. It may be preferred to carry out padding at a temperature below 20°C, for example 0 to 10°C.

The degree of substitution of the cellulose fibres achieved is preferably at least 0.15 carboxymethyl group per glucose unit, and is most preferably in the range 0.2 to 0.5. Higher degrees of substitution than 0.5 carboxymethyl group per glucose unit may lead to fibres which are water-soluble rather than water-swellable. Higher degrees of substitution within the above range may be preferred when the fibre is to be used in absorbent products, and lower degrees of substitution within this range may be preferred when the fibre is to be used in garments.

presence of polyvalent cations, particularly polyvalent metal ions, in at least one reagent solution may 20 be advantageous. Preferred metal ions are calcium ions, but barium, magnesium or zinc are alternatives. We have found that the presence of polyvalent metal ions during processing helps to avoid the formation of fibres which are watersoluble in distilled or demineralised water; it is believed 25 that the polyvalent metal ion may be forming cross-links between carboxylic acid groups on different cellulose chains. The concentration of polyvalent metal salt, for example calcium salt, in the treatment solution can for example be in the range 0.01 to 10% by weight. In many 30 cases, calcium ions present in hard tap water may have a sufficient effect. In other cases it may be preferred to add a calcium salt such as calcium chloride to the treatment solution, for example in an amount of 1 to 4% by weight. In general, the higher the degree of substitution of the 35 cellulose fibre with carboxymethyl groups, the higher is the preferred concentration of calcium ions (within the stated

range) in the reagent solution to avoid forming fibres which are soluble in distilled or demineralised water. The polyvalent metal ions, for example calcium ions, can alternatively be applied during washing of the treated fibre by including them in the washing liquid, but this is less effective.

In a modification of the process of the invention, the fibre is first treated with aqueous strong alkali so that the alkali is absorbed throughout the fibre. The fibre is 10 then rinsed with a solvent for the alkali, preferably water, without washing thoroughly. The water may optionally contain one or more surfactants. The effect of such rinsing is to remove more alkali from the outer region of the fibre than fibre is then treated with the from the interior. The 15 monochloroacetic reagent, preferably monochloroacetic acid, and is heated to cause carboxymethylation and drying of the fibre. The level of carboxymethylation is greater at the interior of the fibre where more alkali remains than at the outer regions of the fibre. A rinsing step can be carried 20 out after treatment with the monochloroacetic reagent and before heating and drying; this tends to increase the difference in the degree of carboxymethylation between the interior and the outer regions of the fibre.

The product of this modified process, namely a 25 carboxymethylated cellulose fibre having a higher degree of carboxymethylation at the interior of the fibre than in the outer regions of the fibre, feels similar to conventional cellulosic fibres even when wet but has an increased absorbency.

30 After the carboxymethylation process, the fibre is usually washed to remove any unreacted alkali or chloroacetate or any by-products such as sodium chloride or sodium glycollate. An aqueous wash is generally used, preferably a mixture of water with a water-miscible organic solvent. The washing medium may contain an organic hydroxyl

- 9 -

compound, a surfactant, and/or an acid. The organic hydroxyl compound is a compound containing at least one alcoholic hydroxyl group, for example ethanol, methanol or another low molecular weight alcohol and/or a polyhydroxy compound such 5 as ethylene glycol or propylene glycol. The weight ratio of the organic hydroxyl compound to water can for example be in the range 3:1 to 1:50. A low molecular weight mono-alcohol can act both as water-miscible organic solvent and as organic hydroxyl compound; for example a preferred washing 10 medium is based on a mixture of water and ethanol in weight ratio 2:1 to 1:2. If a surfactant is used it is preferably a non-ionic surfactant such as a polyalkylene oxide adduct of an alcohol or phenol, although anionic or cationic surfactants can be used. Any surfactant used should 15 preferably be hydrophilic rather than hydrophobic; such a hydrophobic surfactant may reduce the rate of water uptake by the fibres. Examples of preferred surfactants are those sold under the Trade Marks "Tween 20" and "Atlas G1086". An acid used during washing to neutralise the alkalinity of the 20 carboxymethylated fibre is preferably a weak acid, for example an organic carboxylic acid such as acetic acid which is used for example at 0.5 to 15% by weight, preferably 1 to 5%. The weight ratio of wash liquor to fibre used during washing is preferably in the range 5:1 to 50:1.

washing is preferably carried out by a counter-current washing procedure, for example in 2 or 3 stages. In a 2-stage counter-current washing procedure, fibre which has already been washed once is washed again with clean wash liquor. The washed fibre from this second wash stage can be dried for further processing. The liquor resulting from this second wash stage passes to the first wash stage as the wash liquor for unwashed fibre. Counter-current washing allows the use of a lower wash liquor to fibre weight ratio, for example a ratio of 10:1 can be used to wash as effectively as a ratio of 20:1 used in a single washing step. A relatively low concentration of acid, for example 0.5 to 2.0% by weight, can be used in the wash liquor.

- 10 -

As an alternative to inclusion of a surfactant in the wash liquor, it may be preferred to apply a surfactant subsequently as a finish. The surfactant can for example be applied as a solution in an alcohol or an aqueous alcohol mixture, for example the mixture used to wash the fibre, or a liquid surfactant can be applied undiluted. The finish can be applied by immersion of the fibre in the finish, or by lick roller or by spray. If the surfactant is applied as a finish, the fibre is preferably pressed to remove excess 10 wash liquor, for example by mangling, before applying the finish.

After the required washes, the fibre is generally dried. Excess wash liquor is preferably removed by pressure, for example by mangling, followed by heat drying. The optimum degree to which the fibre should be dried depends on the intended further processing, but a moisture content of 5-20% by weight is generally preferred to avoid brittleness of the dried fibres, particularly when drying a tow, yarn or staple fibre which has to undergo further processing such as 20 crimping, carding, weaving or felting.

The treated fibre can be crimped, and it may be preferred to crimp the fibre to give increased loft, particularly if the fibre is to be used in non-woven applications for absorbent products. A tow can for example be crimped by stuffer box crimping. False twist crimping can alternatively be used. If the fibre is to be crimped, it may be possible to omit the heat drying step after washing and to allow the fibre to become dried during crimping.

Carboxymethylated fibre produced from solvent-spun cellulose according to the invention has higher absorbency and superior physical properties compared to carboxymethylated fibre produced from regenerated cellulose or cotton fibres. The absorbency of 0.9% saline solution, as measured by the free-swell method, can for example be 15 or 35 more, e.g. 20 to 40, grams per gram, combined with a

- 11 -

tenacity in the range 25-15cN/tex. Viscose rayon or cotton fibres carboxymethylated by the same process absorbencies only in the range 8-13 g/g and a lower tenacity. At this level of absorbency, carboxymethylated 5 viscose rayon in particular and carboxymethylated cotton to a lesser extent become sticky on the surface in contact with moisture, so that when a tow of fibres is carboxymethylated the fibres become glued together and lose their individual fibrous nature. This problem is not encountered with the 10 fibres of the present invention, which can be processed using conventional textile machinery, for example by the staple route including cutting, carding and if desired crimping, drafting and spinning. Even at lower degrees of substitution giving lower absorbency, the carboxymethylated 15 solvent-spun cellulose fibres are substantially stronger than carboxymethylated viscose rayon fibres.

The form of the carboxymethylated fibres after swelling in water or in an aqueous liquid such as saline solution depends on the absorbency of the fibres and the diameter of 20 the fibres. Absorbency generally increases with increasing carboxymethyl group content. At high levels of absorbency the swollen fibres tend to form a coherent gel in which the identity of individual fibres is lost, particularly if the fibres are of low decitex. For example, fibres of initial 25 decitex 1.7 per filament and having an absorbency (0.9% saline solution, free swell) of 28 g/g, corresponding to treatment with 19.2% by weight ClCH2COONa and 6.5% NaOH, swell to a gel in tap water of hardness 400 p.p.m. CaCO₃. Fibres of the same initial decitex, treated with 13.3% 30 ClCH $_2$ COONa and 4.5% NaOH and having an absorbency of 20, remain as discrete fibres when swollen in tap water and can be re-dried to fibrous form. Fibres of initial decitex 6.0, treated with 22.1% C1CH2COONa and 7.5% NaOH and having a absorbency of 27, also remain as discrete fibres when 35 swollen in tap water. For absorbent disposable products, gel formation on swelling is acceptable. For non-disposable garments, retention of fibrous form when wet and re-dried is

- 12 -

necessary.

The carboxymethylated fibre of the present invention can be used in various products. It can, for example, be used in absorbent personal products such as tampons, 5 disposable diapers, sanitary napkins or incontinence pads. carboxymethylated fibres are preferably used one or more other textile fibres, combination with preferably consisting wholly or mainly of cellulosic fibres, for example cellulosic fibres such as cotton or regenerated 10 cellulose or fibres having a higher absorbency than most textile fibres but less than carboxymethyl cellulose fibres, such as multi-limbed cellulose fibres as described in EP-Aare preferably fibres carboxymethylated 301874. intimately mixed with said other fibres, for example by 15 carding or air-laying the fibres together to form a web of mixed fibres, or they can be used as a layer, for example a non-woven fabric, of carboxymethylated fibres sandwiched between layers of said other fibres. The proportion of carboxymethylated fibres in a blend with cellulosic fibres 20 for absorbent products can for example be at least 5% up to 95%, preferably at least 10% and up to 50%, The carboxymethylated particularly 15-25%, by weight. fibres can also be used at similar levels in conjunction products. absorbent wood pulp in fluffed with 25 Carboxymethylated fibres can be used alone in absorbent personal products, particularly those having a relatively low degree of substitution and relatively low absorbency, but it is preferred to use a blend of carboxymethylated fibres having a relatively high degree of substitution and 30 absorbency with non-carboxymethylated cellulose fibres. The invention, carboxymethylated cellulose fibre of the particularly that having a relatively low degree of substitution, can be used in garments such as underwear or sportswear to give increased absorbency and comfort. For 35 such uses, the carboxymethyl cellulose fibre is usually blended with another fibre, preferably a cellulose fibre such as viscose rayon, including multi-limbed viscose, or

- 13 -

cotton, but alternatively a synthetic fibre such as polypropylene or polyester.

The fibre described above having a higher degree of carboxymethylation at the interior of the fibre than in the outer regions is also suitable for use in garments such as underwear or sports wear. It is less effective than uniformly carboxymethylated fibre in absorbent products since its rate of uptake of aqueous fluid is lower, as is its overall absorbency.

needled or hydroentangled fabric formed of solvent-spun cellulose fibre can be carboxymethylated to form a fabric which swells on contact with water to form a liquid-proof barrier and which can be used for example for wrapping electrical components. A yarn or fabric tape of the carboxymethyl cellulose fibre of the invention can be used to wrap cable or can be laid longitudinally in the cable to prevent water ingress.

The carboxymethyl cellulose fibre of the invention can 20 be used as an absorbent for materials such as camphor or menthol or for perfumes, for example in devices adapted to give slow release of these materials. Slow release of camphor or menthol may be desired for medical uses. Slow release of perfume may be desired in air-freshening devices.

The carboxymethyl cellulose fibre of the invention having a relatively low degree of substitution can be used in papermaking, helping to form strong bonds between fibres on drying.

The carboxymethyl cellulose fibre can be used as an 30 absorbent fibre in many other uses, for example in filters, in absorbent liners or mats for packaging, disposable wipes, shoe insoles, swellable gaskets or seals, moisture retention mats in horticulture, moisture-retaining packaging or

- 14 -

swellable self-sealing stitching threads.

The invention is illustrated by the following Examples, in which parts and percentages are by weight:-

Example 1

5

Sodium hydroxide and sodium monochloroacetate were separately dissolved in water and the solutions were cooled to 18°C. The solutions were mixed to form a treatment solution containing 8.5% NaOH and 22.1% ClCH2COONa. A tow of 1.7 decitex solvent-spun cellulosic fibres, the fibres having a substantially uniform structure across their cross-section, as sold under the Trade Mark "Tencel", was padded with the treatment solution for 2 minutes and mangled to give a pick-up (increase in weight of wet fibres) of about 150%. The treated tow was dried at 180°C for 4 minutes, during which time reaction of the cellulose of the fibres to form sodium carboxymethyl cellulose took place. The dried tow was washed with a solution containing 50% ethanol (industrial alcohol), 35% water, 5% glycerol and 10% acetic 20 acid, and was re-dried.

The tow produced had a degree of substitution in excess of 0.1, a tenacity of 24.8 cN/tex and an extensibility of 15.5%. It absorbed 39 grams tap water (hardness 460 p.p.m. CaCO₃) per gram of fibre when allowed to swell freely. The 25 fibres dissolved at least partially in distilled or demineralised water and could not be used as an absorbent for these purified waters; in practical use the fluids being absorbed are generally tap water or aqueous fluids richer in minerals than tap water. The absorbency of the tow for 0.9% saline solution would be somewhat less than for tap water but would be above 20 grams per gram of fibre.

The free swell absorbency was measured by dispersing 0.5 g fibre in 30 ml water and leaving it for 5 minutes. For all absorbency measurements, the fibre is conditioned at 65%

- 15 -

relative humidity and 20°C before being tested. The dispersion is then filtered through a sintered Mark 1 funnel of pore size 100-160 microns and is left for 5 minutes, or until it stops dripping. The water filtered through the funnel is weighed and the weight of water absorbed by the fibres is calculated by subtraction.

Examples 2 and 3

The procedure of Example 1 was repeated using different times and temperatures for drying the treated tow (reaction 10 times and temperatures). In both cases the degree of substitution was above 0.1 and the tenacity above 10 cN/tex. The results were as follows:-

	Example No	Temperature of Drying	Time of Drying	Free Swell Absorbency in
15				Tap Water
10	2	80°C	10 minutes	44 g/g
	3	57°C	26 minutes	28 g/g

Examples 4 and 5

The procedure of Example 1 was repeated using different 20 concentrations of reagents in the treatment solution. In both cases the degree of substitution was above 0.1. The results obtained were as follows:-

	Example No	Concen	tration of	Tenacity	Extensibility	Absorbe	ncy
0.5		I	Reagent	(cN/tex)			tap
25		NaOH	ClCH2COONa	(2011)		wa	ter
	4 .	4.0%	10.4%	30.6	11.8%	8.7	g/g
	5	6.5%	16.9%	24.9	11.3%	25	g/g
30	Untreated fibres	-	-	38.0	14.0%	3.5	g/g

(The free swell absorbency of the product of Example 4 in 0.9% saline solution was about 8 g/g.)

- 16 -

Examples 6 and 7

The procedure of Example 1 was repeated except that calcium chloride was added to the tap water used to prepare the solutions of sodium hydroxide and sodium 5 monochloroacetate. The concentration of calcium chloride was 2% (Example 6) and 3% (Example 7). The free swell absorbency in tap water of the tow produced was 33 g/g and 28 g/g respectively. The tenacity was above 20 cN/tex and the degree of substitution above 0.1 in both cases.

10 Example 8

A tow of 1.7 decitex solvent-spun cellulosic fibres was padded with an 8.5% aqueous solution of NaOH to give a pick-up of about 250%. The treated tow was dried at 180°C for 8 minutes. The tow was then immersed in a 22.1% aqueous solution of ClCH2COONa at 80°C for 30 minutes. The tow was washed and dried as described in Example 1. The properties of the fibres produced were similar to those of the fibres of Example 2.

Example 9

A tow of 1.7 decitex solvent-spun cellulosic fibres was padded with a 22.1% aqueous solution of ClCH₂COONa and mangled to give a pick-up of about 150%. The treated tow was dried at 180°C for 4 minutes. The tow was then padded with an 8.5% aqueous solution of NaOH and mangled to give a pick-up of about 150%. The treated tow was again dried at 180°C for 4 minutes. The dried tow was washed and dried as described in Example 1.

Example 10

The process of Example 9 was repeated except that the 30 order of treatment was reversed; the tow was first treated with sodium hydroxide and then with sodium

- 17 -

monochloroacetate.

The fibres produced in Example 9 and in Example 10 each had properties similar to the fibres of Example 1.

Example 11

A tow of Tencel fibres having a filament decitex of 1.7 was obtained in a never-dried state. The tow was passed through a hand mangle. The amount of water left on the tow 3after mangling was 62%. This wet tow was put in a solution containing 7.5% sodium hydroxide and 22.1% sodium 10 monochloroacetate at room temperature (20°C) for 2 minutes. The padded tow was mangled again. The total pick-up after mangling was 75%. The padded and mangled tow was then reacted in a conditioning cabinet set at 23% RH (relative humidity) and 90°C for five minutes. The amount of water 15 retained on the tow after the treatment was 13%.

After heat treatment the tow was washed in a solution containing 55% industrial alcohol, 42% water and 3% acetic acid. Washed tow was then treated with a finish containing 99% industrial alcohol and 1% Atlas G1086 emulsifier. After 20 this, the tow was dried at a low temperature, leaving some residual moisture on the fibres. The finished tow was crimped using a stuffer box system. The crimped fibres were cut into staple.

The fibres had a degree of substitution above 0.1, a 25 tenacity of 22.5 cN/tex and an extensibility of 12%. The moisture regain of fully dried fibres at 65% RH was 17%. The free swell absorbency of the fibres was measured by the process of Example 1 but using 0.9% saline solution in place of water. The absorbency was 30 g/g. The retention of the 30 saline solution after application of pressure at about 3.4 kPa for 5 minutes or until dripping stops was also measured by weighing the water expressed after application of pressure. The absorbency retention of the fibres was 20 g/g.

Comparative Examples

- A. The procedure of Example 11 was repeated using a tow of "Fibro" regenerated cellulose (viscose rayon) of the same filament decitex. The carboxymethylated viscose rayon fibres produced had a free swell absorbency in 0.9% saline solution of 11 g/g, with a retention of 9 g/g after application of pressure. The carboxymethylated viscose rayon fibres, unlike the carboxymethylated solvent-spun cellulose fibres, became sticky on the surface after brief contact with water.
- 10 B. The procedure of Example 11 was also repeated using combed cotton yarn. The carboxymethylated cotton had a free swell absorbency of 10.5 g/g and a retention of 9 g/g after application of pressure.

Examples 12 to 15

1.7 was padded with a solution of sodium hydroxide and sodium monochloroacetate. The concentrations of the reagents differed in different Examples as shown below. The tow was lightly mangled to stop dripping and the total pick-up was measured. The tow was then dried at 90°C to a moisture level of 13%.

The resulting tow was washed in a solution containing 55% ethanol, 42% water, 2.5% acetic acid and 0.5% citric acid. The washed tow was treated with a finish and dried as described in Example 11. The free swell absorption of the fibres in 0.9% saline solution was measured, as was the retention under 3.42 kPa. The degree of substitution (number of carboxymethyl groups per glucose unit) was also measured. The results were as follows:-

- 19 -

	Example No.	of Rea	tration gents % .CH ₂ COONa	Pick-up	Degree of Substitution	Free Ret Swell Absorb- ency g/g	ention g/g	State of Swollen Fibre
5	12	4.5	13.3	200	0.235	20	10	Fibrous
_	13	5.5	16.2	230	0.29	18	10	Fibrous
	14	6.5	19.2	230	0.375	28	18	Gel
	15	7.5	22.1	275	0.405	38	29	Gel

10 The tenacity of the fibres reduced with increasing degree of substitution but was in all cases greater than 15cN/tex.

Examples 16 and 17

The process of Example 15 was repeated using samples of solvent-spun cellulosic fibre tows of different 15 filament decitex. The results obtained were as follows:

	Example No.	Initial filament decitex		Tenacity cN/tex	bility	Free Ret Swell Absorb- ency g/g	ention g/g	State of swollen fibre
20	16	3.0	265	25.6	17.7	31	22	Gel
	17	6.0	273	18.8	17.7	27	17	Fibrous

Examples 18 to 20

15% carboxymethylated fibres produced according to the 25 invention were carded in a blend with 85% multi-limbed regenerated cellulose fibres, sold under the trade mark "Galaxy", using a Shirley Miniature Card. The carboxymethylated fibres used were:

- Example 18 crimped fibres produced according to Example 30 11
 - Example 19 fibres produced according to Example 11 omitting the crimping stage
 - Example 20 crimped fibres produced according to Example 15.

PCT/GB92/02283

- 20 -

The carded webs were each formed into a radially expanding tampon and tested for absorbency in a "modified Syngina" test as defined in GB-A-2094637, pages 4 to 6, except that a 200 mm hydrostatic head air pressure was 5 employed. The absorbency in the modified Syngina test was tested with a 1% saline solution. The results obtained were:

Example 18	6.5	g/g
Example 19	6.0	g/g
Example 20	7.6	g/g

10 By comparison, the absorbency of a similar tampon formed from a 100% "Galaxy" carded web in this test is 5.1 g/g.

Examples 21 and 22

The fibres produced in Example 15 were blended with 15 fluffed wood pulp (a mixture of the pulps sold under the Trade Marks "Sappi" and "Caima" for use in diapers) in different ratios. Such blends of wood pulp and absorbent material are used commercially in light-weight diapers. The absorbency of the blends in terms of free swell in 0.9% 20 saline solution and retention after application of pressure was measured. As a comparison, sodium carboxymethyl cellulose (CMC) powder sold under the Trade Mark "Courlose" (having the highest absorption level amongst commercial CMC powders) was blended with the same wood pulp in the same 25 ratios. The results obtained were:

	Example No.	Ratio	ç	of C	MC fibre	B Free Swell	Retention	Comparative using CMC	Results- powder
						Absorbency g/g	g/g	Free Swell g/g	Retention g/g
30	21	15	5	:	85	22	12	20	12
50	22	50	0	:	50	34	25	23	16
	_	C	0	:	100	12	8		

- 21 -

As shown by these results, the carboxymethyl cellulose fibres according to the invention increased absorbency by a similar amount to the CMC powder when used at the 15% level, but provided a much greater absorbency when used at the 50% 5 level.

Example 23

A solution of 6.5% sodium hydroxide and 19.2% sodium monochloroacetate was prepared and cooled to -2°C in a treatment bath. A tow of never-dried "Tencel" 1.7 decitex 10 filaments was passed at 5m/min successively through a roller nip of 100 kPa (to reduce the water content to 62% based on dry tow), the above treatment bath, a roller nip of 34 kPa (to give a total solution pick-up of 75%) and a drying cabinet at 90°C/10% RH for 7 minutes. The treated tow was 15 washed as described in Example 11 and was re-dried and cut into staple.

The fibre produced had a degree of substitution above 0.1, a free swell absorbency in 0.9% saline solution of 34.1 g/g with a retention of 24.4 g/g, and a tenacity above 10 cN/tex. Fibre of this absorbency could be produced continuously if the temperature of the treatment bath was maintained at around 0°C, but the absorbency properties of the fibre produced fell slightly over time if the temperature of the treatment bath was allowed to rise.

CLAIMS

- 1. Carboxymethyl cellulose fibre having a degree of substitution of at least 0.1 carboxymethyl group per glucose unit, characterised in that the fibre is derived from solvent-spun cellulose fibre and has an absorbency of at least 8 grams 0.9% saline solution per gram of fibre, as measured by the free swell method, and a tenacity of at least 10cN/tex.
- Carboxymethyl cellulose fibre according to claim 1,
 characterised in that it has a degree of substitution of
 to 0.5 carboxymethyl group per glucose unit.
- 3. Carboxymethyl cellulose fibre according to claim 1 or claim 2, characterised in that it has an absorbency of 15 to 40 grams 0.9% saline solution per gram of fibre, as 15 measured by the free swell method.
 - 4. Carboxymethyl cellulose fibre according to any of claims 1 to 3, characterised in that it has a tenacity of 15 to 25cN/tex.
- 5. Carboxymethyl cellulose fibre according to any of 20 claims 1 to 4, characterised in that it is capable of remaining as discrete fibres when swollen with water of hardness 400 p.p.m. CaCO₃.
- 6. A process for the production of carboxymethyl cellulose fibre by reacting cellulose fibre with a strong 25 al ili and a monochloroacetic reagent selected from monochloroacetic acid and salts thereof, characterised in that the cellulose fibre used is solvent-spun cellulose fibre.
- 7. A process according to claim 6, characterised in 30 that the solvent-spun cellulose fibre is reacted when in tow

- 23 -

or staple fibre form.

•

8. A process according to claim 6 or claim 7, characterised in that the solvent-spun cellulose fibre used is never-dried fibre.

- 9. A process according to claim 6, characterised in that the solvent-spun cellulose fibre is reacted in the form of a woven, knitted or non-woven fabric.
- 10. A process according to any of claims 6 to 9, characterised in that the solvent-spun cellulose fibre is 10 contacted with an aqueous solution containing 4 to 10% by weight alkali metal hydroxide and 10 to 25% by weight monochloroacetate salt.
- 11. A process according to any of claims 6 to 10, characterised in that the solvent-spun cellulose fibre is 15 heated to a temperature in the range 80 to 150°C after the strong alkali and the monochloroacetic reagent have been applied to the fibre.
- 12. A process according to claim 11, characterised in that the strong alkali and the monochloroacetic reagent are 20 applied to the solvent-spun cellulose fibre together from an aqueous solution which is at a temperature of 20°C or below.
- 13. A process according to claim 11, characterised in that separate solutions of alkali metal hydroxide and of monochloroacetate salt are applied to the solvent-spun 25 cellulose fibre before it is heated.
- 14. An absorbent personal product comprising a layer which is a mixture of carboxymethyl cellulose fibre and one or more other textile fibres, characterised in that the said layer comprises 10 to 50% by weight carboxymethyl cellulose 30 fibre according to any of claims 1 to 5 or produced by a process according to any of claims 6 to 13.

- 24 -

15. An absorbent personal product according to claim 14, characterised in that the other textile fibres consist wholly or mainly of cellulosic fibres.

16. An absorbent personal product comprising a layer 5 which is a mixture of absorbent fibre and fluffed wood pulp, characterised in that the absorbent fibre is carboxymethyl cellulose fibre according to any of claims 1 to 5 or produced by a process according to any of claims 6 to 13, the carboxymethyl cellulose fibre being present in the layer 10 at 10 to 50% by weight.

INTERNATIONAL SEARCH REPORT

International Application No

PCT/GB 92/02283

I. CLASS	SIFICATION OF SUBJECT MATTER (il sever	at classification symbols apply, indicate all)						
According	According to International Patent Classification (IPC) or to both National Classification and IPC IPC ⁵ : D 01 F 11/02, D 01 F 2/28, A 61 L 15/28							
IL FIELDS	S SEARCHED	Documentation Searched 7						
Classification		Classification Symbols						
	•							
IPC ⁵	D 01 F,A 61 L,A 6							
	Documentation Searche to the Extent that such Do	d other than Minimum Documentation currents are included in the Fields Searched ⁶						
III. DOCU	MENTS CONSIDERED TO BE RELEVANT	•	Relevant to Claim No. 13					
Category *		here appropriate, of the relevant passages 12	Referent to Ciefat No.					
			•					
x	DE, A1, 3 209 126 (KAO SOAP CO.,	LTD.)	1,2,6,					
	28 October 1992 claims; page 3	, lines 6-13;						
	page 11, table	1.						
	& GB, A, 2 094 (cited in the a	application).						
x	us, A, 4 044 766		1,2,6,					
^	(KACZMARZYK et	al.)	14,15					
,	30 August 1977 claims; column	2. line 60 -						
	column 3, line	11.						
x	GB, A, 2 220 881		1,2,6					
	(TOYO BOSEKI K	ABUSHIKI						
	KAISHA) 24 Jani (24.01.90),	nary 1990						
İ	claims: page 2	, lines 30-33.						
	(cited in the	application).						
	US, A, 4 246 221	- -	1,6					
A	U3, A, 4 240 221	·						
	I categories of cited documents: 10 ument defining the general state of the ert which	"I" leter document published after or priority date and not in cont cited to understand the princi- invention	ple or theory underlying the					
con: "E" earli	sidered to be or particular resevance for document but published on or after the intermediates.	stional exe document of particular releva						
"L" doc	ument which mey throw doubts on priority claim this cited to establish the publication dete of an iden or other special reason (as specified)		nce; the claimed invention an inventive step when the					
-0- doc	ument referring to an oral disclosure, use, exhibit	tion or document is combined with on ments, such combination being	obvious to a person skilled					
202 does	ument published prior to the international filing de r than the priority date claimed	te but "4" document member of the same	petent family					
IV. CERT	FICATION	Date of Melling of this International	Search Report					
Date of the	Actual Completion of the International Search	2.6 M	AR 1993					
	15 March 1993	Signature of Authorized Officer						
Internation	al Searching Authority	WEIGERSTORFE	R e.h.					
	EUROPEAN PATENT OFFICE WEIGHTON EN E.H.							

	ENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEE	Relevant to Claim No.
*800A .	Chation of Document, ** with indication, where appropriate, of the relevant passages	
	(McCORSLEY, III)	1
	20 January 1981 (20.01.81),	
ĺ	claims.	
	(cited in the application).	
- 1		
1		
į		
1.		
1		
- 1	•	:
İ		
l	•	
İ		
l		
- 1	•	
- 1		
- 1		
1		
	•	
.		
- 1		
l l		
	İ	
1		
		
1		
f	·	
1		
ļ		
	j	
1		
-		
Ī		
1		

zum internationalen Recherchenbericht über die internationale Patentanmeldung Nr. to the International Search Report to the International Patent Application No.

au rapport de recherche international relatif à la demande de brevet international n°

PCT/GB 92/02283 SAE 67825

angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Unter-richtung und erfolgen ohne Gewähr.

In diesem Anhang sind die Mitglieder der Patentfamilien der im obenge-nannten internationalen Recherchenbericht angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Unterrichtung und erfolgen ohne Gewähr.

This Annex lists the patent family members relating to the patent documents members de la famille de brevets relatifs aux documents de brevets dans le rapport de recherche international visée ci-dessus. Les resments fournis sont donnés à titre of information.

relatifs aux documents de brevets cités dans le rapport de recherche inter-national visée ci-dessus. Les reseigne-ments fournis sont donnés à titre indica-tif et n'engagent pas la responsibilité de l'Office.

			05 1 0111771 ·					
angeführte Patent in sea Document	erchembericht s Patentdokument document cited rch report de brevet cité pport de recherche	Datum der Veröffentlichung Publication date Date de publication	Mitglied(er) der Patentfamilie Patent family member(s) Membre(s) de la famille de brevets	Datum der Veröffentlichung Publication date Date de publication				
DE A1	3209126	28-10-82	DE C2 3209126 FR A1 2502191 FR B1 2502191 GB A1 2094802 GB B2 2094802 JP A2 57153001 JP B4 60015641 US A 4410694	18-10-90 24-09-82 18-04-86 22-09-82 20-02-85 21-09-82 20-04-85 18-10-83				
US A	4044766	30-08-77	AU A1 22204/77 AU B2 498198 BR A 7701153 CA A1 1045801 DE A1 2709132 FR A1 2342054 GB A 1535135 IT A 1086210 JP A2 52105694 NL A 7702032 ZA A 7701149	17-08-78 15-02-79 18-10-77 09-01-79 08-09-77 23-09-77 06-12-78 28-05-85 05-09-77 30-08-77 25-01-78				
GB A	2220881	·	GB A1 2220881 GB B2 2220881 JP A2 1280069	24-01-90 08-07-92 10-11-89				
US A	4246221	20-01-81	048085555887077744009585558877077744400958887370777444009695888330972216677922499966932167767678442944797722449946687370777445772224999669321677666787335599667835598833222499466879722249479722466990064797429446987722249944695991118447747686888688888888888888888888888	150-101-1-1001-1011-100114-100115715-1012314-1001231239392008599039946099931 2010-101-1-1-10011-1-1001				

3.

- - -- -

=

۴